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Please find below and/or attached an Office communication concerning this application or proceeding.

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	Application No.	Applicant(s)	
	09/492,954	PYLE ET AL.	
Office Action Summary	Examiner	Art Unit	-
	Jeanine A. Goldberg	1634	
The MAILING DATE of this communication ap	ppears on the cover sheet	with the correspondence address	
A SHORTENED STATUTORY PERIOD FOR REPI WHICHEVER IS LONGER, FROM THE MAILING I - Extensions of time may be available under the provisions of 37 CFR 1 after SIX (6) MONTHS from the mailing date of this communication. - If NO period for reply is specified above, the maximum statutory period - Failure to reply within the set or extended period for reply will, by statu Any reply received by the Office later than three months after the maili earned patent term adjustment. See 37 CFR 1.704(b).	DATE OF THIS COMMUNITY .136(a). In no event, however, may divil apply and will expire SIX (6) More, cause the application to become	NICATION. a reply be timely filed ONTHS from the mailing date of this communicatio ABANDONED (35 U.S.C. § 133).	
Status			
1)⊠ Responsive to communication(s) filed on 29 . 2a)⊠ This action is FINAL . 2b)□ Th 3)□ Since this application is in condition for allows closed in accordance with the practice under	is action is non-final. ' ance except for formal m	· ·	s
Disposition of Claims			
4) ☐ Claim(s) 1 and 6-8 is/are pending in the appli 4a) Of the above claim(s) is/are withdress 5) ☐ Claim(s) is/are allowed. 6) ☐ Claim(s) 1 and 6-8 is/are rejected. 7) ☐ Claim(s) is/are objected to. 8) ☐ Claim(s) are subject to restriction and/	awn from consideration.		
Application Papers			
9) The specification is objected to by the Examination The drawing(s) filed on is/are: a) acceptable and applicant may not request that any objection to the Replacement drawing sheet(s) including the correct 11) The oath or declaration is objected to by the Examination is objected to by the Examination The specification The specif	ccepted or b) objected to objected to objected to object of objection is required if the drawi	rance. See 37 CFR 1.85(a). ng(s) is objected to. See 37 CFR 1.121((d).
Priority under 35 U.S.C. § 119		•	
12) Acknowledgment is made of a claim for foreig a) All b) Some * c) None of: 1. Certified copies of the priority documer 2. Certified copies of the priority documer 3. Copies of the certified copies of the pri application from the International Bure: * See the attached detailed Office action for a list	nts have been received. nts have been received in ority documents have be au (PCT Rule 17.2(a)).	Application No en received in this National Stage	
Attachment(s) 1) \(\sum \) Notice of References Cited (PTO-892)	4) ☐ Intervie	v Summary (PTO-413)	
 2) Notice of Draftsperson's Patent Drawing Review (PTO-948) 3) Information Disclosure Statement(s) (PTO-1449 or PTO/SB/08 Paper No(s)/Mail Date 	Paper N	o(s)/Mail Date f Informal Patent Application (PTO-152)	

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DETAILED ACTION

This action is in response to the papers filed August 29, 2005. Currently, claims
 6-8 are pending.

- 2. All arguments have been thoroughly reviewed but are deemed non-persuasive for the reasons which follow.
- 3. Any objections and rejections not reiterated below are hereby <u>withdrawn</u> in view of the amendments to the claims.
- 4. This action contains new grounds of rejection necessitated by amendment.
- 5. This action is FINAL.

Information Disclosure Statement

6. The listing of references in the specification is not a proper information disclosure statement. 37 CFR 1.98(b) requires a list of all patents, publications, or other information submitted for consideration by the Office, and MPEP § 609 A(1) states, "the list may not be incorporated into the specification but must be submitted in a separate paper." Therefore, unless the references have been cited by the examiner on form PTO-892, they have not been considered.

Response

It is noted that an IDS was filed on January 27, 2000. The IDS was considered on June 22, 2000 and mailed to applicant. There are numerous references on pages 41-44 of the specification which are not listed on this 1449. Thus, as stated previously,

unless the references have been cited by the examiner on form PTO-892, they have not been considered.

Claim Rejections - 35 USC § 103

The following is a quotation of 35 U.S.C. 103(a) which forms the basis for all obviousness rejections set forth in this Office action:

- (a) A patent may not be obtained though the invention is not identically disclosed or described as set forth in section 102 of this title, if the differences between the subject matter sought to be patented and the prior art are such that the subject matter as a whole would have been obvious at the time the invention was made to a person having ordinary skill in the art to which said subject matter pertains. Patentability shall not be negatived by the manner in which the invention was made.
- 7. This application currently names joint inventors. In considering patentability of the claims under 35 U.S.C. 103(a), the examiner presumes that the subject matter of the various claims was commonly owned at the time any inventions covered therein were made absent any evidence to the contrary. Applicant is advised of the obligation under 37 CFR 1.56 to point out the inventor and invention dates of each claim that was not commonly owned at the time a later invention was made in order for the examiner to consider the applicability of 35 U.S.C. 103(c) and potential 35 U.S.C. 102(e), (f) or (g) prior art under 35 U.S.C. 103(a).
- 8. Claims 1 and 7-8 are rejected under 35 U.S.C. 103 (a) over Shuman (Proc. Natl. Acad. Sci. USA, November 1992, Vol. 89, pages 10935-10939) in view of Bjornson et al. (Biochemistry, (1994). Vol. 33, pages 14306-14316) as <u>evidenced</u> by Stern et al. (US Pat. 5,712,096, January 1998) and Karn et al. (US Pat. 6,316,194, Nov. 2001).

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Shuman teaches a method for detecting the release of a single-stranded RNA from an RNA duplex which comprise admixing an RNA helicase with the RNA duplex under conditions permitting the RNA duplex to unwind the RNA duplex and release single stranded RNA, wherein the RNA duplex comprises a first RNA having a label and a second RNA wherein the unwound single-stranded RNA released from the duplex is detected by gel electrophoresis (Page 10936, Col. 1, lines 18-29, and 40-52, and Figures 1-2). Shuman teaches a method, wherein ATP and a divalent cation is present (Methods Section. Enzyme Assays Subsection). The enzyme assays used helicase reaction mixtures which contained Tris-HCL, 2mM dithiothreitol, mgCl2, GTP, 20-50 fmol of P-labeled standard dsRNA substrate (20,000,000-50,000,000 nM). Shuman teaches a method of measuring the rate of release of a single-stranded RNA from an RNA duplex which complies detecting whether the single-stranded RNA is released from the RNA duplex at predetermined time intervals, and detecting therefrom the rate of release of the single-stranded RNA from the RNA duplex (Results Section, Kinetics Subsection and Figure 2). Shuman teaches a method of determining whether a compound is capable of modulating the release of a single-stranded RNA from an RNA duplex (Results Section, Requirements of Helicase Activity Subsection).

Shuman does not teach the method, wherein the first label is capable of producing a luminescent energy pattern wherein the first RNA is present in the RNA duplex which differs from the luminescent energy pattern produced when the first RNA is not present in the RNA duplex, thereby detecting release of a single-stranded RNA from the RNA duplex.

Shuman does not teach the method. wherein the first label is present at the 5' end of the first RNA and the second label is attached to the 3' end of the second RNA and the luminescent energy pattern results from interaction of luminescent energy released from the first label with the second label.

However Bjornson et al. teach the method, wherein the first label is capable of producing a luminescent energy pattern wherein the first nucleotide is present in the nucleic acid duplex which differs from the luminescent energy pattern produced when the first nucleotide is not present in the nucleotide duplex, thereby detecting release of a single-stranded nucleic acid from the nucleic acid duplex after admixing helicase (Abstract. and Results section). Bjornson teaches that the fluorescence assay is extremely sensitive allowing DNA unwinding reactions to be monitored continuously at DNA concentrations as low as **1nM** in a fluorescence stopped-flow experiment (abstract). Moreover, Bjornson et al.teach several advantages of using a fluorescent based assay for kinetic studies in general and particularly for mechanistic studies for helicase-catalyzed unwinding. Bjornson et al. teach the method, wherein the first label is present at the 5' end of the first nucleic acid and the second label is attached to the 3' end of the second nucleic acid and the luminescent energy pattern results from interaction of luminescent energy released from the first label with the second label. Materials and Methods Section, Preparation of DNA unwinding subsection, and Results section and Figure 1).

Further, Stern teaches labeling both the 5' and 3' end of RNA. Stern teaches the RNA termini with fluorescein phosphoroamidites (3' labeling) or CPG (5' lablelin) or the

incorporation of fluorescent adenosine or cytosine nucleotides at specific positions internal in the RNA (col. 15, lines 15-20).

Moreover, Karn teaches FRET labeleling of RNA. Karn teaches that the target RNA may be fluorescently labeled at the 3' or 5' end of a strand within the target RNA or within the chain of the target RNA. Karn teaches an entire section of fluorescent labeling which includes labeling with 2 fluorescent groups with one group laced adjacent to the 5' end of the target RNA and a second fluorescent group placed adjacent to the 3' end of the target (col. 11-12).

Therefore, it would have been prima facie obvious to one of ordinary skill at the time the invention was made to have substituted and combined the method of Shuman in view of Bjornson. The ordinary artisan would have been motivated to have substituted the fluorescent labeling method of Bjornson for the radiolabeled method of Shuman. The ordinary artisan would have been motivated to have the first label that is capable of producing a luminescent energy pattern wherein the first nucleotide is present in the nucleic acid duplex which differs from the luminescent energy pattern produced when the first nucleotide is not present in the nucleotide duplex, thereby detecting release of a single-stranded nucleic acid from the duplex after admixing helicase of Bjornson et al in the method of Schuman. The art clearly teaches that ability to label both the 5' and 3' end of RNA as well as DNA. Bjornson specifically states, "we describe a fluorescence assay that can be used to monitor helicase-catalyzed unwinding of duplex DNA continuously in real time (Abstract, first sentence). The

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ordinary artisan would have been motivated to have used a continuously real time assay in lieu of a radiolabeled method without the functionality of real time detection. Further motivation is provided by Bjornson, "this emphasizes the utility of the continuous spectroscopic method described here, which allows many more time points to be collected, thus enabling more accurate determinations of the complete time course and observed rate constants for all phase of a multiphasic reaction (Page 1431 6, Column 1, last sentence of the second paragraph). An ordinary artisan would have been motivated to have substituted and combined the method, wherein the first label is capable of producing a luminescent energy pattern wherein the first nucleotide is present in the nucleic acid duplex which differs from the luminescent energy pattern produced when the first nucleotide is not present in the nucleotide duplex, thereby detecting release of a single-stranded nucleic acid from the duplex after admixing helicase of Bjomson in the method of Schuman, in order to achieve the express advantages, as noted by Bjornson, of a fluorescence assay that can be used to monitor helicase-catalyzed unwinding of duplex nucleic acids continuously in real time and which emphasizes the utility of the continuous spectroscopic method described here allowing many more time points to be collected, thus enabling more accurate determinations of the complete time course and observed rate constants for all phase of a multiphasic reaction.

Response to Arguments

The response traverses the rejection. The response asserts that "none of the cited references teach the detection of nanomolar amounts of fluorescently labeled

RNA." It is noted that not one of the references individually teaches detection of nanomolar amounts of fluorescently labeled RNA. However, given the teachings in the art, the ordinary artisan would have been motivated to have detected RNA at low concentrations and would have had a reasonable expectation of success as exemplified by Bjornson's ability to detect low concentrations of DNA. Bjornson teaches detecting DNA helicase activity using 1nM and fluorescently labeled duplex DNA. The art teaches that RNA may be fluorescently labeled on both the 5' and 3' end. Thus, performing the RNA method of Shuman by substituting the fluorescent method of Bjornson in view of the state of the art would provide a method for detecting the release of ssRNA from RNA duplex using fluorescence. As provided above, the ordinary artisan would have been motivated to have used a continuously real time assay in lieu of a radiolabeled method without the functionality of real time detection. Further motivation is provided by Bjornson, "this emphasizes the utility of the continuous spectroscopic method described here, which allows many more time points to be collected, thus enabling more accurate determinations of the complete time course and observed rate constants for all phase of a multiphasic reaction (Page 1431 6, Column 1, last sentence of the second paragraph). Bjornson teaches the fluorescence assay is extremely sensitive allowing DNA unwinding reactions to be monitored continuously at DNA concentrations as low as 1nM which is well within any defined nanomolar range. The art teaches analyzing the unwinding of both DNA and RNA. The art teaches analyzing very low quantities of DNA. The ordinary artisan would have a reasonable expectation that a detection method for RNA at low concentrations would be successful since the art

teaches both RNA and DNA unwinding may be detected. Therefore, the art suggest that the fluorescent assay for detecting unwinding may be monitored using a concentration of nucleic acid as low as 1nM. Given the teachings of Shuman in view of Bjornson, there would have been an reasonable expectation of success for detecting RNA in the nanomolar range using a RNA helicase assay.

Thus for the reasons above and those already of record, the rejection is maintained.

9. Claim 6 is rejected under 35 U.S.C. 103 (a) over Shuman (Proc. Natl. Acad. Sci. USA, November 1992, Vol. 89, pages 10935-10939) in view of Bjornson et al. (Biochemistry, (1994), Vol. 33, pages 14306-14316) further in view of Nazarenko et al. (US Pat. 5,866,336, February 1999).

Neither Shuman nor Bjornson teach the labels fluorescein isothiocyanate and rhodamine isothiocyanate.

However, Nazarenko et al. (herein referred to as Nazarenko) teaches an extensive list of suitable moieties that can be selected as donor or acceptors in FRET pairs (col. 17-18).

It would have been prima facie obvious to a practitioner having ordinary skill in the art at the time the invention was made to have substituted and combined the labels fluorescein isothiocyanate and rhodamine isothiocyanate of Nazarenko in the method of Schuman in view of Bjornson. Bjornson teaches using fluorescein and hexachlorofuorescein which are among the listed donors and acceptors. Therefore,

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using alternative donors and/or acceptors which were known in the art would have the ability to quench signals as the labels taught by Bjornson. Therefore, using equivalent labels in the method would have been obvious to the ordinary artisan. An ordinary artisan would have been motivated to substitute and combine the labels fluorescein isothiocyanatc and rhodamine isothiocyanate of Nazarenko in the method of Schuman in view or Bjornson because Nazarenko teaches that the FRET donors and acceptors are functional equivalents.

Response to Arguments

The response traverses the rejection. The response asserts Nazarenko does nothing to overcome the deficiencies of Shuman or Bjornson in failing to teach the detection of nanomolar amounts of fluorescently labeled RNA. This argument has been reviewed but is not convincing because the combination of teachings of Shuman or Bjornson render obvious the detection of nanomolar amounts of fluorescently labeled RNA for the reasons set forth above. Thus for the reasons above and those already of record, the rejection is maintained.

10. Claims 1, 7-8 are rejected under 35 U.S.C. 103(a) as being unpatentable over Eggleston (Nucleic Acids Research, Vol. 24, No. 7, pages 1179-1186, 1996).

Eggleston et al. (herein referred to as Eggleston) teaches a helicase assay based upon the displacement of fluorescent nucleic acid binding ligands. The helicase assay is continuous, kinetic assay based on the displacement of the fluorescent dyes from dsDNA upon DNA unwinding. Eggleston analyzes several dyes including ethidium

bromide to function as suitable reporter molecules. Eggleston teaches that a variety of fluorophores were examined to determine their utility as reporter molecules in a continuous helicase assay. Reactions using several of these dyes share the properties of having relatively low fluorescence in the presence of ssDNA and significant fluorescence enhancement upon binding to dsDNA (page 1180, col. 2). Eggleston teaches that they anticipate that this dye displacement assay can find widespread use in the study of RNA helicases, thereby suggesting a reasonable expectation of success. Eggleston suggests that the dye displacement assay can be readily adapted for use with other DNA helicases, with RNA helicases and with other enzymes that act on nucleic acids.

Eggleston teaches the details of the fluorometric helicase assay and how the assay was performed, measured and optimized depending upon the dye and enzyme concentration used (page 1181, col. 1). For the fluorometric helicase assay, the DNA substrate was 10 uM (0.01 nM) nucleotides BamHI-digested pBR322." (page 1181, col.

- 1). The unwinding was initiated by the addition of ATP in excess of Mg2+ ion concentrations (limitations of Claim 8). As seen in the Figure 1, the nucleic acid is labeled at the 5' end. It is noted that the 5' end is not the 5' terminus. The 5' end is interpreted to mean nucleotides 5' of the middle of the nucleic acid (limitations of Claim
- 1). The assay is performed over a relative time, for example at 10 time intervals, see Figure 1 (limitations of Claim 7).

Eggleston teaches that H33258 displays the greatest dsDNA specificity relative to ssDNA followed by TO, EB and DAPI. Due to their specificity for dsDNA, strong

fluorescence signal, and minimal fluorescence in the absence of DNA, the fluorophores DAPI, H33258 and TO were selected for further study.

Eggleston states that "the studies have focused on DNA helicases, but the dye displacement assay may provide a new means by which the unwinding activity of RNA helicases can be examined" (page 1185, col. 2). Eggleston teaches that "since this dye binds to RNA in addition to DNA, it is readily conceivable that RNA helicases may be amenable to this assay if an appropriate ligand, such as EB or perhaps, propidium iodide, is utilized" (page 1185, col. 2). Eggleston teaches that "once this parameter is optimized for a particular substrate and enzyme, the dye-displacement assay gives results which are consistent with those obtained from other types of assays" (page 1185). Therefore, while Eggleston's studies focus primarily on DNA helicases, Eggleston specifically teaches that the method would be applicable to RNA helicases and there would be a reasonable expectation of success for the RNA helicases method. Therefore, it would have been prima facie obvious to one of ordinary skill at the time the invention was made to have performed the dye-displacement assay using RNA helicase to analyze and study the continuous unwinding of RNA using the method taught by Eggleston. With regard to the specific amount of RNA duplex present between 1-3 nm, the ordinary artisan would have recognized that the ability to detect 0.01nm would enable the detection of a greater amount of RNA duplex, including 1-3 nanomolar. Therefore detecting a greater amount of RNA would have been obvious depending on the amount of RNA duplex present.

Response to Arguments

The response traverses the rejection. The response asserts Eggleston fails to teach the element of using between 1-3 nanomolar. This argument has been thoroughly reviewed, but is not found persuasive. As addressed in the rejection above, "With regard to the specific amount of RNA duplex present between 1-3 nm, the ordinary artisan would have recognized that the ability to detect 0.01nm would enable the detection of a greater amount of RNA duplex, including 1-3 nanomolar. Therefore detecting a greater amount of RNA would have been obvious depending on the amount of RNA duplex present." Detecting greater amounts of RNA would have a reasonable expectation of success.

The response asserts that the ordinary artisan, even if the dye was reasonably expected to bind to RNA, would not have necessarily expected the claimed method to succeed absent applicants' own experimentation. This argument has been thoroughly reviewed, but is not found persuasive because the response does not provide arguments particular to why the ordinary artisan would not have expected the claimed method to succeed.

The response previously asserted that a skilled artisan would recognized that the claimed invention overcomes difficulties not present with DNA, i.e., making an RNA strand and attaching a luminescent dye to it. This argument has been thoroughly reviewed, but is not found persuasive the prior art teaches methods for attaching a luminescent dye to RNA (see Stern and Karn above). Therefore, there would be a reasonable expectation of success for making an RNA strand and attaching a luminescent dye to it.

Thus for the reasons above and those already of record, the rejection is maintained.

11. Claims 1, 7-8 are rejected under 35 U.S.C. 103(a) as being unpatentable over Kowalczykowski et al. (US Pat. 5,747,247, May 1998).

Kowalczykowski et al. (herein referred to as Kowalczykowski) teaches a helicase assay based upon the displacement of fluorescent nucleic acid binding ligands. The helicase assay is continuous, kinetic assay based on the displacement of the fluorescent dyes from dsDNA upon DNA unwinding. Kowalczykowski analyzes several dyes including ethidium bromide to function as suitable reporter molecules. Kowalczykowski teaches that a variety of fluorophores were examined to determine their utility as reporter molecules in a continuous helicase assay. Reactions using several of these dyes share the properties of having relatively low fluorescence in the presence of ssDNA and significant fluorescence enhancement upon binding to dsDNA (col 3, lines 30-45). Kowalczykowski teaches that they anticipate that this dye displacement assay can find widespread use in the study of RNA helicases, thereby suggesting a reasonable expectation of success. Kowalczykowski suggests that the dye displacement assay can be readily adapted for use with other DNA helicases, with RNA helicases and with other enzymes that act on nucleic acids.

Kowalczykowski teaches the details of the fluorometric helicase assay and how the assay was performed, measured and optimized depending upon the dye and enzyme concentration used (col 7-8). For the fluorometric helicase assay, the DNA

substrate was 10 uM (0.01nM) nucleotides BamHI-digested pBR322 which is within "the nanomolar range." The unwinding was initiated by the addition of ATP in excess of Mg2+ ion concentrations (limitations of Claim 2, 8). As seen in the Figure 1, the nucleic acid is labeled at the 5' end. It is noted that the 5' end is not the 5' terminus. The 5' end is interpreted to mean nucleotides 5' of the middle of the nucleic acid (limitations of Claim 3). The assay is performed over a relative time, for example at 10 time intervals, see Figure 1 (limitations of Claim 7).

Kowalczykowski teaches that H33258 displays the greatest dsDNA specificity relative to ssDNA followed by TO, EB and DAPI. Due to their specificity for dsDNA, strong fluorescence signal, and minimal fluorescence in the absence of DNA, the fluorophores DAPI, H33258 and TO were selected for further study.

Kowalczykowski states that "the dye displacement assay also provides a new means by which the unwinding activity of RNA helicases can be examined (col. 15, lines 4-5). Kowalczykowski teaches that "since this dye binds to RNA in addition to DNA, RNA helicases are likewise amenable to this assay if an appropriate ligand, such as EB or perhaps, propidium iodide, is utilized" (col. 15, lines 13-15). Kowalczykowski teaches that "the dye displacement assay can be adapted for use with any helicase, whether it utilizes a DNA or RNA substrate, provided that a suitable, minimally-inhibitory nucleic acid binding dye is selected" (col. 15, lines 15-20). Therefore, while Kowalczykowski's studies focus primarily on DNA helicases, Kowalczykowski specifically teaches that the method would be applicable to RNA helicases and there would be a reasonable expectation of success for the RNA helicases method.

Therefore, it would have been prima facie obvious to one of ordinary skill at the time the invention was made to have performed the dye-displacement assay using RNA helicase to analyze and study the continuous unwinding of RNA using the method taught by Kowalczykowski.

Response to Arguments

The response traverses the rejection. The response asserts Kowalczykowski fails to render obvious the claimed invention because Kowalczykowski teaches using .01nM DNA substrate. With regard to the specific amount of RNA duplex present between 1-3 nm, the ordinary artisan would have recognized that the ability to detect 0.01nm would enable the detection of a greater amount of RNA duplex, including 1-3 nanomolar. Therefore detecting a greater amount of RNA would have been obvious depending on the amount of RNA duplex present. Detecting greater amounts of RNA would have a reasonable expectation of success. Thus for the reasons above and those already of record, the rejection is maintained.

Double Patenting

The nonstatutory double patenting rejection is based on a judicially created doctrine grounded in public policy (a policy reflected in the statute) so as to prevent the unjustified or improper timewise extension of the "right to exclude" granted by a patent and to prevent possible harassment by multiple assignees. See *In re Goodman*, 11 F.3d 1046, 29 USPQ2d 2010 (Fed. Cir. 1993); *In re Longi*, 759 F.2d 887, 225 USPQ 645 (Fed. Cir. 1985); *In re Van Ornum*, 686 F.2d 937, 214 USPQ 761 (CCPA 1982); *In re Vogel*, 422 F.2d 438, 164 USPQ 619 (CCPA 1970);and, *In re Thorington*, 418 F.2d 528, 163 USPQ 644 (CCPA 1969).

A timely filed terminal disclaimer in compliance with 37 CFR 1.321(c) may be used to overcome an actual or provisional rejection based on a nonstatutory double patenting ground provided the conflicting application or patent is shown to be commonly owned with this application. See 37 CFR 1.130(b).

Effective January 1, 1994, a registered attorney or agent of record may sign a terminal disclaimer. A terminal disclaimer signed by the assignee must fully comply with 37 CFR 3.73(b).

12. Claims 1, 6-8 are provisionally rejected under the judicially created doctrine of obviousness-type double patenting as being unpatentable over claims 1-8 of copending Application No. 10/182,362 in view of Bjornson et al.

This is a <u>provisional</u> obviousness-type double patenting rejection.

An obviousness-type double patenting rejection is appropriate where the conflicting claims are not identical, but an examined application claim is not patentably distinct from the reference claim(s) because the examined claim is either anticipated by or would have been obvious over, the reference claim(s). See e.g., *In re Berg*, 140 F.3d 1428, 46 USPQ2d 1226 (fed. Cir. 1998); *In re Goodman*, 11 F.3d 1046, 29 USPQ2d 2010 (Fed. Cir. 1993); *In re Longi*, 759 F.2d 887, 225 USPQ 645 (Fed. Cir. 1985).

Here, Claim 1-8 of copending Application No. 10/182,362 recites a method of detecting the release of single stranded RNA from RNA duplex with fluorescence assay. The method of copending Application No. 10/182,362 differs from the claims herein in that it fails to disclose detecting RNA duplex within "the nanomolar range." However, Bjornson teaches a helicase DNA unwinding assay which measures by fluorescence energy transfer which uses concentrations of nucleic acid as low as 1 nM. While the claims differ in RNA and DNA, there would be a reasonable expectation of success based upon the state of the art and the ability to label both RNA and DNA on both the 3' and 5' ends that RNA concentrations as low as 1 nM would similarly have a reasonable

expectation of success. Therefore, it would have been obvious to modify the method of Claims 1-8 of copending Application No. 10/182,362 such that the method would use RNA duplex in an amount within the nanomolar range.

Response to Arguments

The response previously indicated that applicant will consider filing a terminal disclaimer of copending Application No. 10/182,362 when the rejection is no longer provisional.

Conclusion

- 13. No claims allowable over the art.
- 14. Applicant's amendment necessitated the new ground(s) of rejection presented in this Office action. Accordingly, **THIS ACTION IS MADE FINAL**. See MPEP § 706.07(a). Applicant is reminded of the extension of time policy as set forth in 37 CFR 1.136(a).

A shortened statutory period for reply to this final action is set to expire THREE MONTHS from the mailing date of this action. In the event a first reply is filed within TWO MONTHS of the mailing date of this final action and the advisory action is not mailed until after the end of the THREE-MONTH shortened statutory period, then the shortened statutory period will expire on the date the advisory action is mailed, and any extension fee pursuant to 37 CFR 1.136(a) will be calculated from the mailing date of the advisory action. In no event, however, will the statutory period for reply expire later than SIX MONTHS from the date of this final action.

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15. Any inquiry concerning this communication or earlier communications from the examiner should be directed to examiner Jeanine Goldberg whose telephone number is (571) 272-0743. The examiner can normally be reached Monday-Friday from 7:00 a.m. to 4:00 p.m.

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If attempts to reach the examiner by telephone are unsuccessful, the examiner's supervisor, Gary Jones, can be reached on (571) 272- 0745.

Information regarding the status of an application may be obtained from the Patent Application Information Retrieval (PAIR) system. Status information for published applications may be obtained from either Private PAIR or Public PAIR. Status information for unpublished applications is available through Private PAIR only. For more information about the PAIR system, see http://pair-direct.uspto.gov. Should you have questions on access to the Private PAIR system, contact the Electronic Business Center (EBC) at 866-217-9197 (toll-free).

The Central Fax Number for official correspondence is (571) 273-8300.

Jeanine Goldberg
Primary Examiner
September 20, 2005